

# Online machine learning with decision trees

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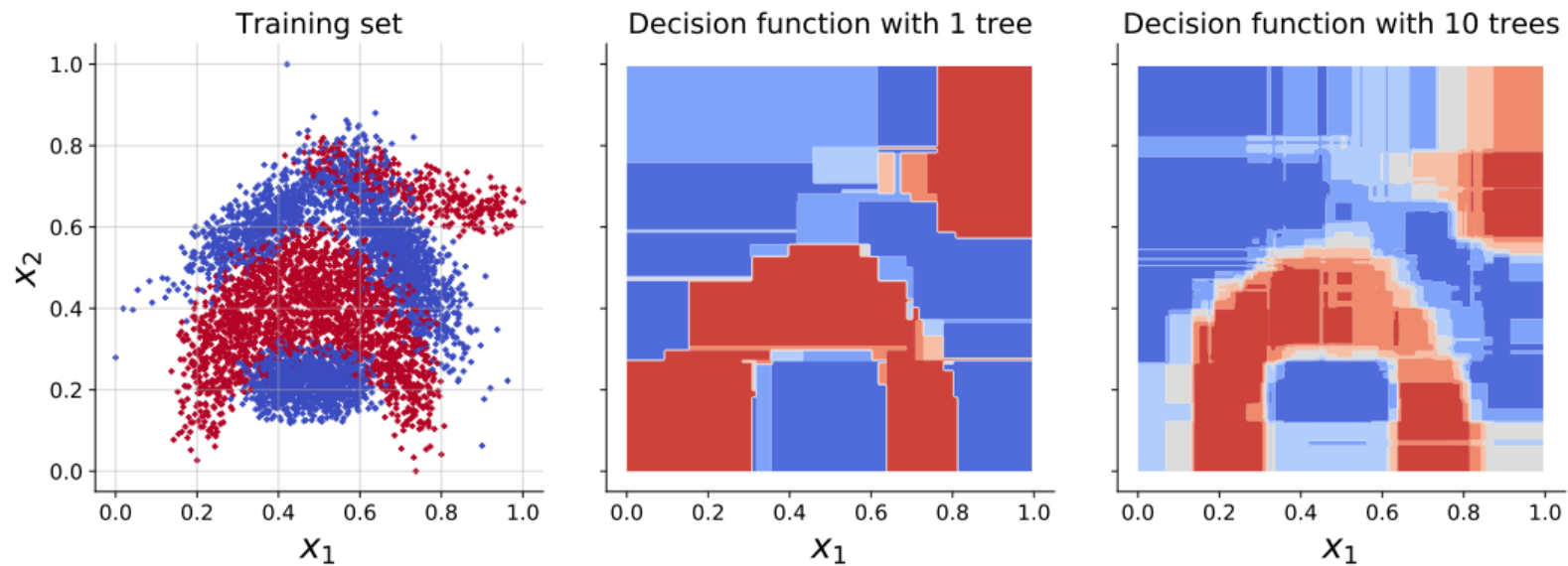
University of Toulouse  
Thursday 7<sup>th</sup> May, 2020

# Decision trees

- “*Most successful general-purpose algorithm in modern times.*” [HB12]
- Sub-divide a feature space into partitions
- Non-parametric and robust to noise
- Allow both numeric and categorical features
- Can be regularised in different ways
- Good weak learners for bagging and boosting [Bre96]
- See [BS16] for a modern review
- Many popular open-source implementations [PVG<sup>+</sup>11, CG16, KMF<sup>+</sup>17, PGV<sup>+</sup>18]

Alas, they assume that the data can be scanned more than once, and thus can't be used in an online context.

# Toy example: the banana dataset<sup>1</sup>



<sup>1</sup>[Banana dataset on OpenML](#)

# Online (supervised) machine learning

- Model learns from samples  $(x, y) \in \mathbb{R}^{n \times p} \times \mathbb{R}^{n \times k}$  which arrive in sequence
- Online != out-of-core:
  - Online: samples are only seen once
  - Out-of-core: samples can be revisited
- **Progressive validation [BKL99]:**  $\hat{y}$  can be obtained right before  $y$  is shown to the model, allowing the training set to also act as a validation set. No need for cross-validation!
- Ideally, **concept drift** [[GŽB<sup>+</sup>14](#)] should be taken into account:
  1. **Virtual drift:**  $P(X)$  changes
  2. **Real drift:**  $P(Y | X)$  changes:
    - ▶ Example: many 0s with sporadic bursts of 1s
    - ▶ Example: a feature's importance changes through time

# Online decision trees

- A decision tree involves enumerating split candidates
- Each split is evaluated by scanning the data
- This can't be done online without storing data
- Two approaches to circumvent this:
  1. Store and update feature distributions
  2. Build the trees without looking at the data (!!)
- Bagging and boosting can be done online [OR01]

# Consistency

- Trees fall under the non-parametric regression framework
- Goal: estimate a regression function  $f(x) = \mathbb{E}(Y | X = x)$
- We estimate  $f$  with an approximation  $f_n$  trained with  $n$  samples
- $f_n$  is consistent if  $\mathbb{E}(f_n(X) - f(X))^2 \rightarrow 0$  as  $n \rightarrow +\infty$
- Ideally, we also want our estimator to be unbiased
- We also want regularisation mechanisms in order to generalise
- Somewhat orthogonal to concept drift handling

# Hoeffding trees

- Split thresholds  $t$  are chosen by minimising an impurity criterion
- The impurity looks at the distribution of  $Y$  in each child
- An impurity criterion depends on  $P(Y | X < t)$
- $P(Y | X < t)$  can be obtained via Bayes' rule:

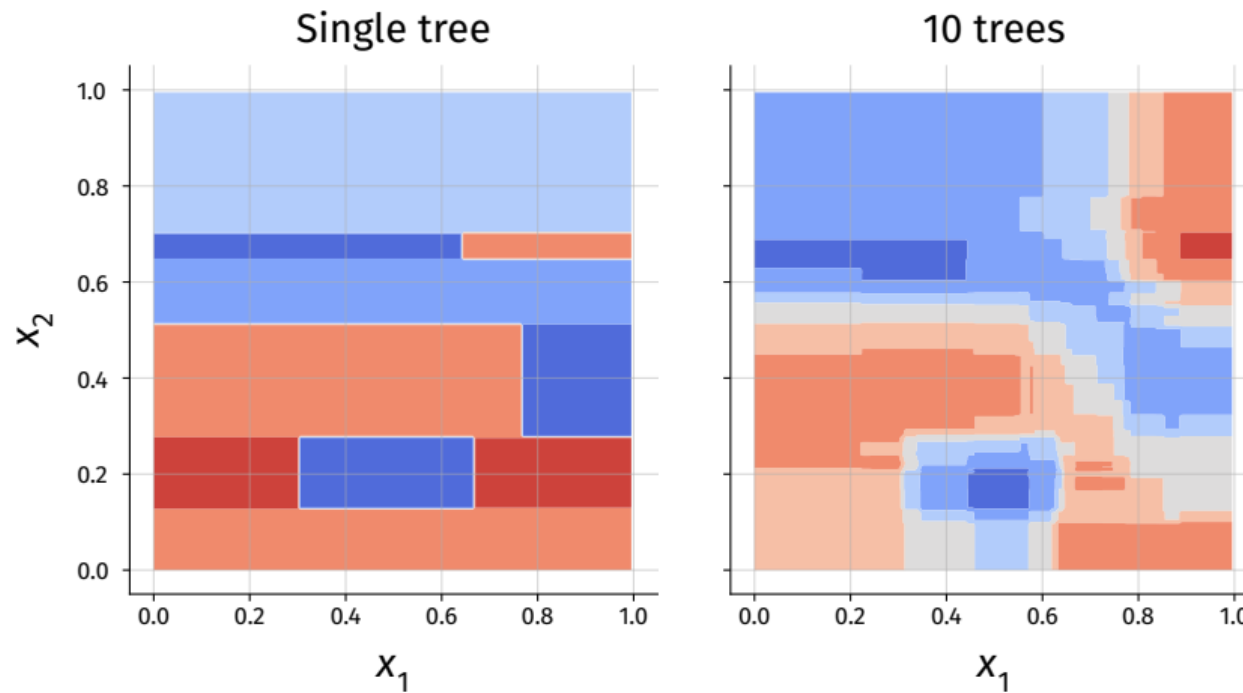
$$P(Y | X < t) = \frac{P(X < t | Y) \times P(Y)}{P(X < t)}$$

- For classification, assuming  $X$  is numeric:
  - $P(Y)$  is a counter
  - $P(X < t)$  can be represented with a histogram
  - $P(X < t | Y)$  can be represented with one histogram per class

# Hoeffding tree construction algorithm

- A Hoeffding tree starts off as a leaf
- $P(Y)$ ,  $P(X < t)$ , and  $P(X < t | Y)$  are updated every time a sample arrives
- Every so often, we enumerate some candidate splits and evaluate them
- The best split is chosen if significantly better than the second best split
- Significance is determined by the Hoeffding bound
- Once a split is chosen, the leaf becomes a branch and the same steps occur within each child
- Introduced in [DH00]
- Many variants, including revisiting split decisions when drift occurs [HSD01]

# Hoeffding trees on the banana dataset



# Mondrian trees

- Construction follows a Mondrian process [RT<sup>+</sup>08]
- Split features and points are chosen without considering their predictive power
- Hierarchical averaging is used to smooth leaf values
- First introduced in [LRT14]
- Improved in [MGS19]

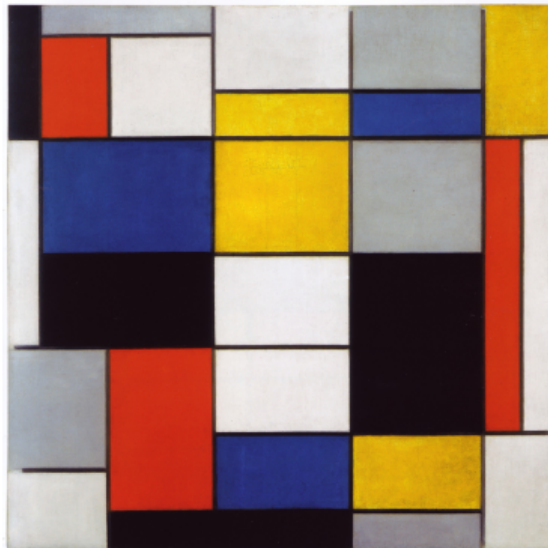
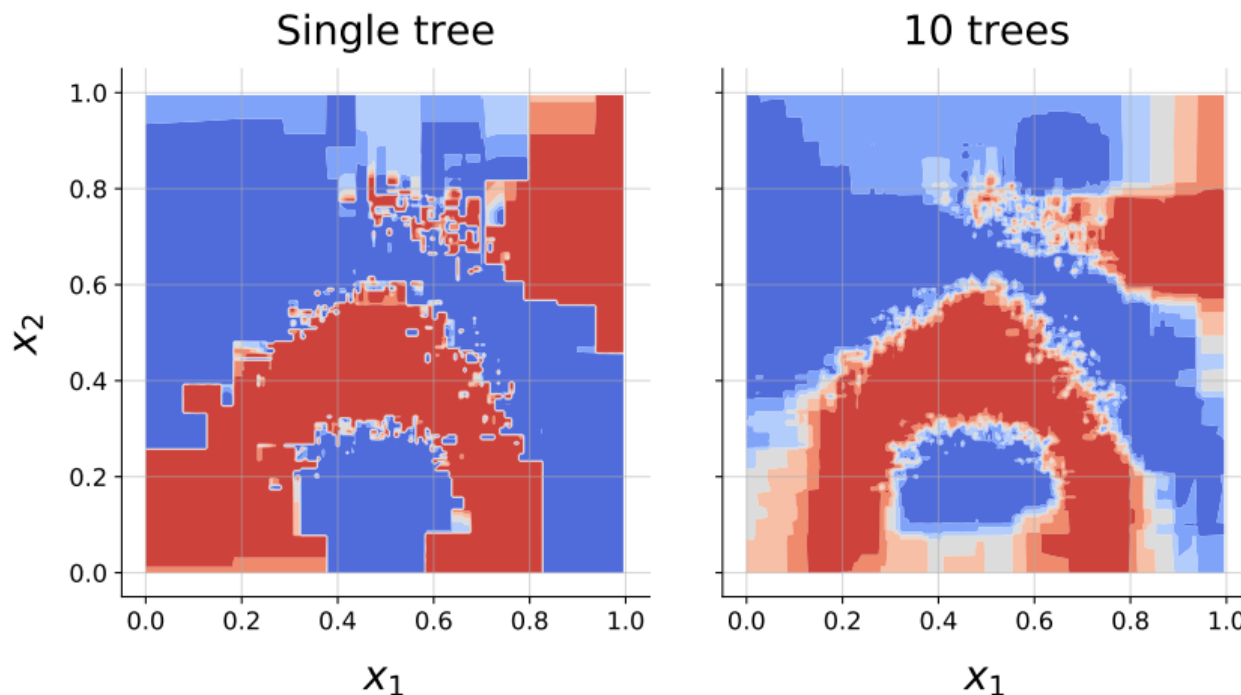


Figure: *Composition A* by Piet Mondrian

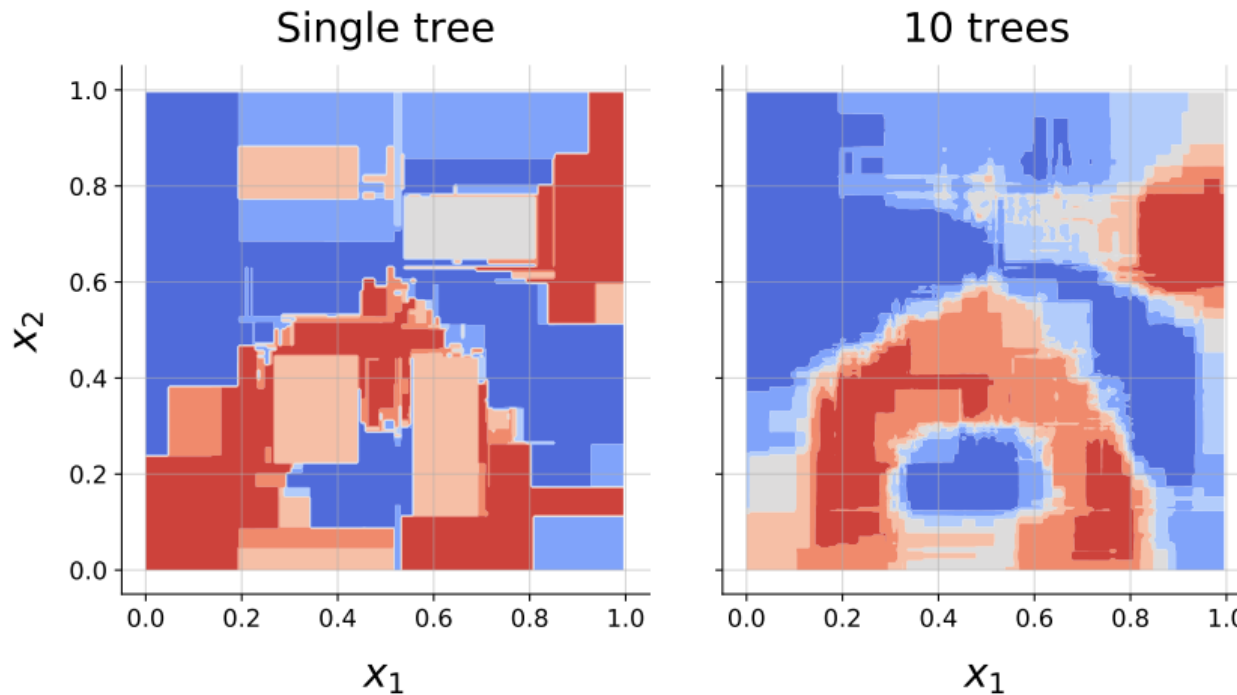
# The Mondrian process

- Let  $u_j$  and  $l_j$  be the bounds of feature  $j$  in a cell
- Sample  $\delta \sim \exp(\sum_{j=1}^p u_j - l_j)$
- Split if  $\delta < \lambda$
- The chances of splitting decrease with the size of the cells
- $\lambda$  is a soft maximum depth parameter
- Features are uniformly chosen in proportion to  $u_j - l_j$
- More information in [these slides](#)

# Mondrian trees on the banana dataset



# Aggregated Mondrian trees on the banana dataset



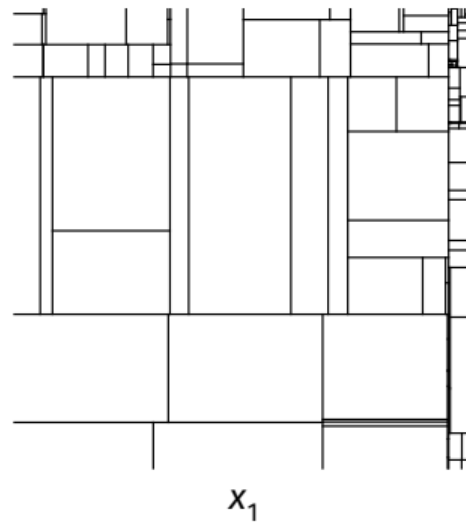
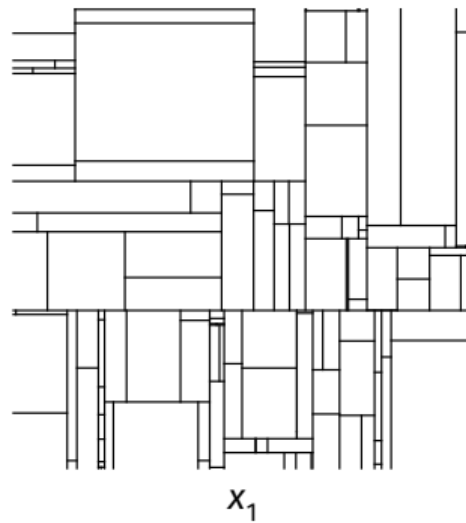
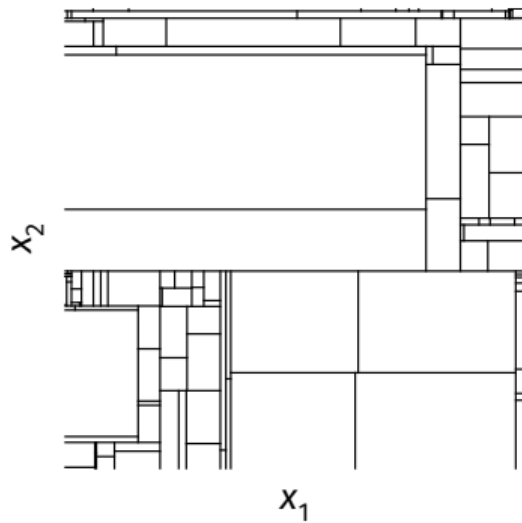
# Purely random trees

- Features  $x$  are assumed to in  $[0, 1]^P$
- Trees are constructed independently from the data, before it even arrives:
  1. Pick a feature at random
  2. Pick a split point at random
  3. Repeat until desired depth is reached
- When a sample reaches a leaf, said leaf's running average is updated
- Easier to analyse because tree structure doesn't depend on  $Y$
- Consistency depends on:
  1. The height of a tree – denoted  $h$
  2. The amount of features that are “relevant”
- Bias analysis performed in [AG14]
- Word of caution: this is different from extremely randomised trees [GEW06]

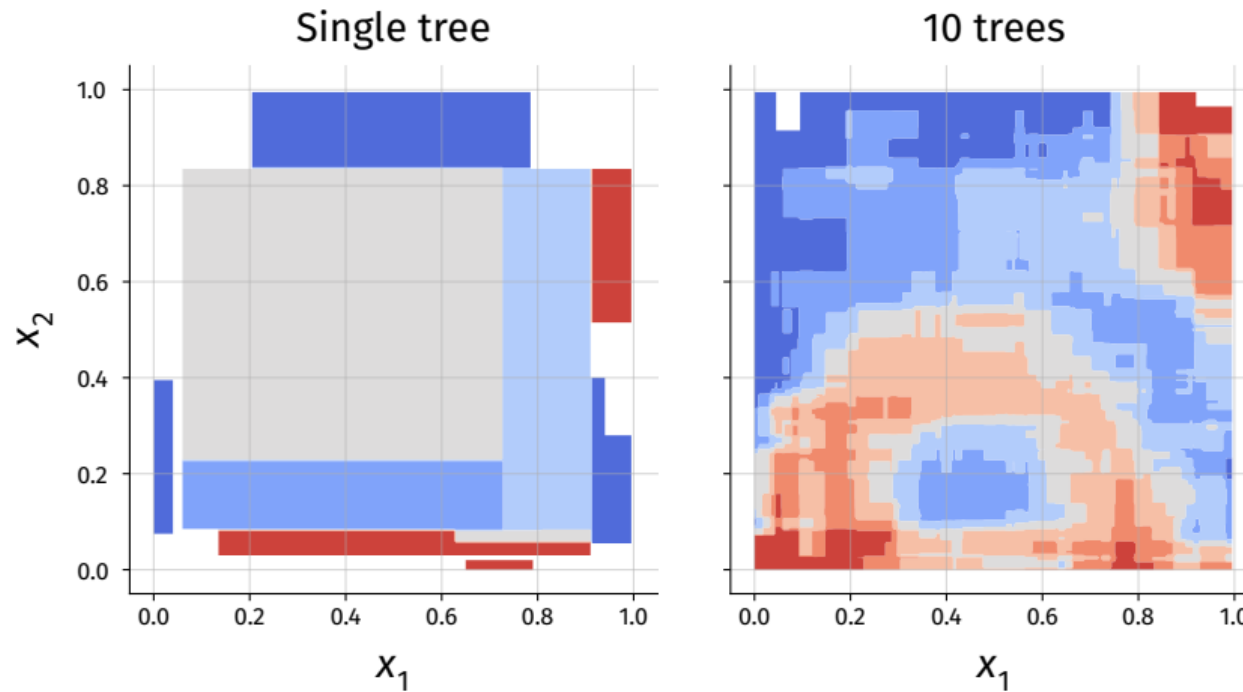
# Uniform random trees

- Features and split points are chosen completely at random
- Let  $h$  be the height of the tree
- Consistent when  $h \rightarrow +\infty$  and  $\frac{h}{n} \rightarrow 0$  as  $h \rightarrow +\infty$  [BDL08]

# Uniform random trees



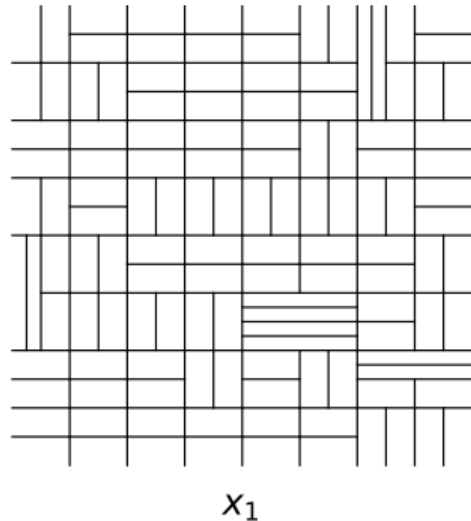
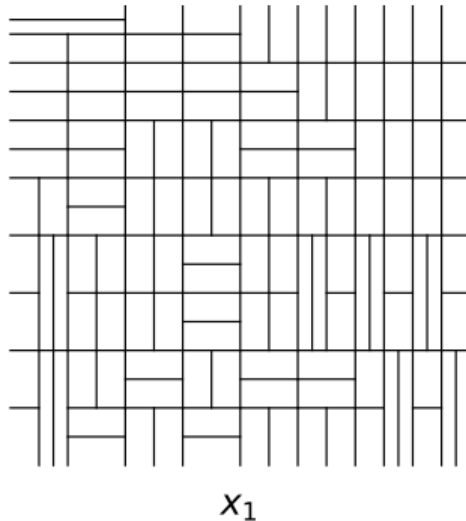
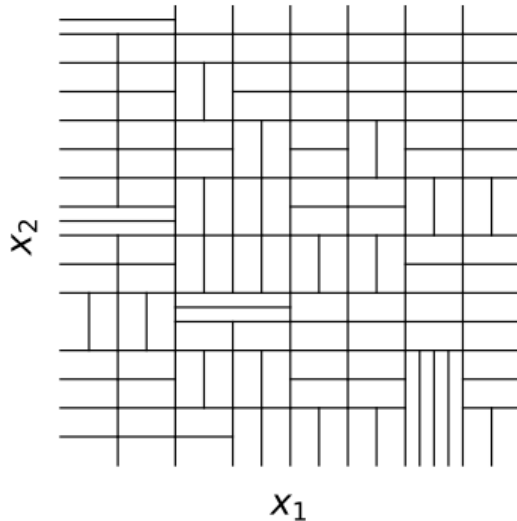
# Uniform random trees on the banana dataset



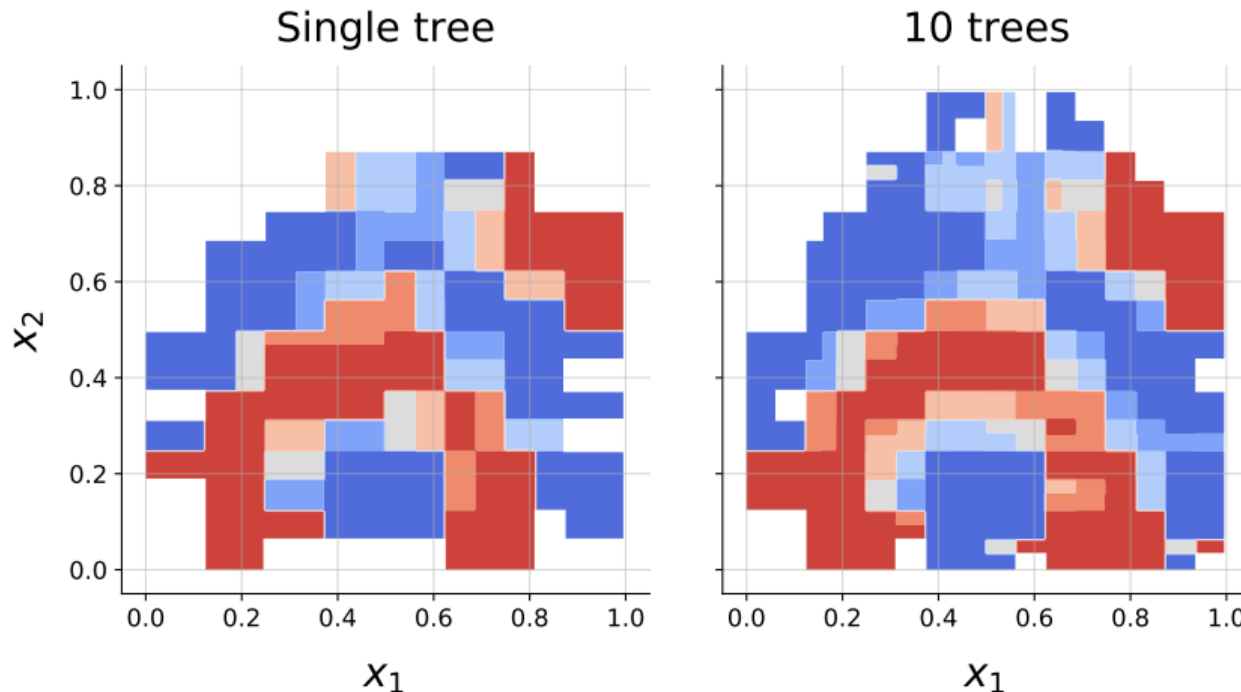
# Centered random trees

- Features are chosen completely at random
- Split points are the mid-points of a feature's current range
- Consistent when  $h \rightarrow +\infty$  and  $\frac{n}{2^h} \rightarrow 0$  as  $h \rightarrow +\infty$  [Sco16]

# Centered random trees

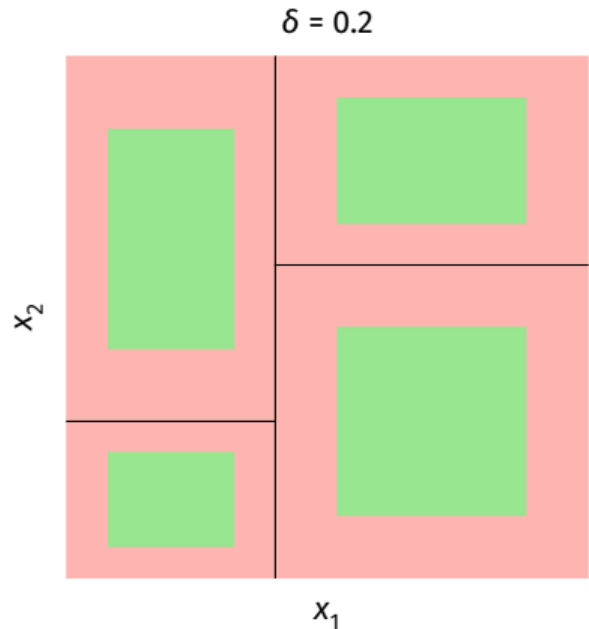


# Centered random trees on the banana dataset

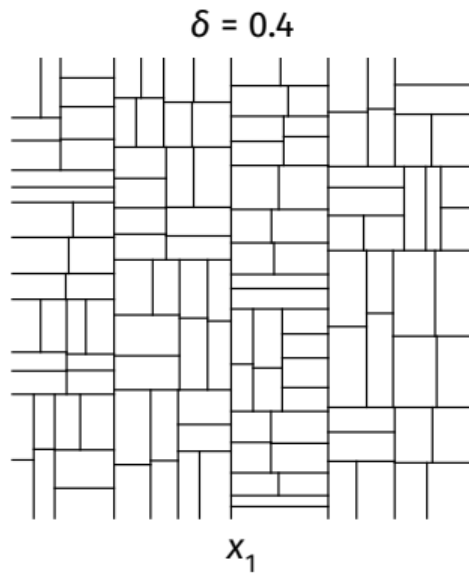
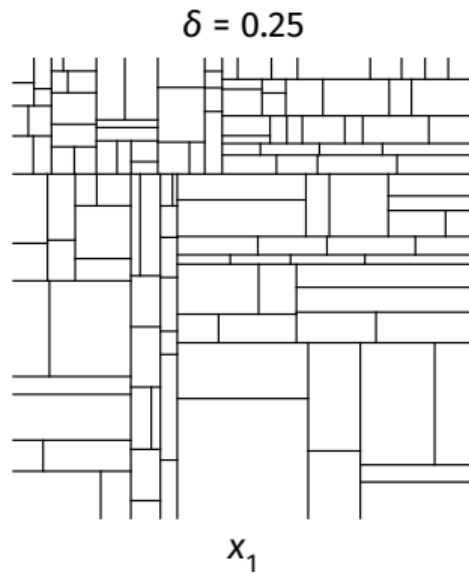
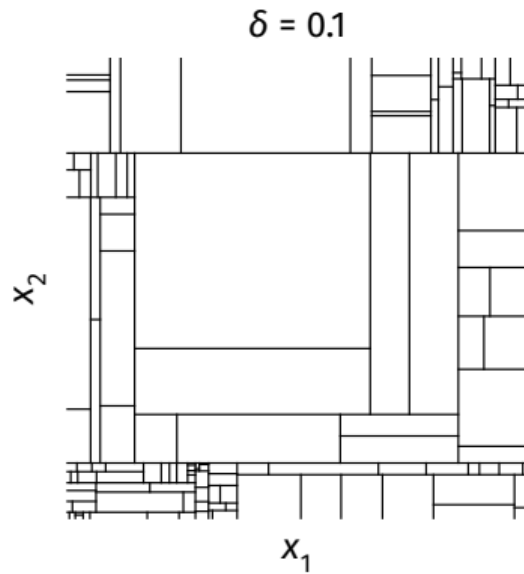


# How about a compromise?

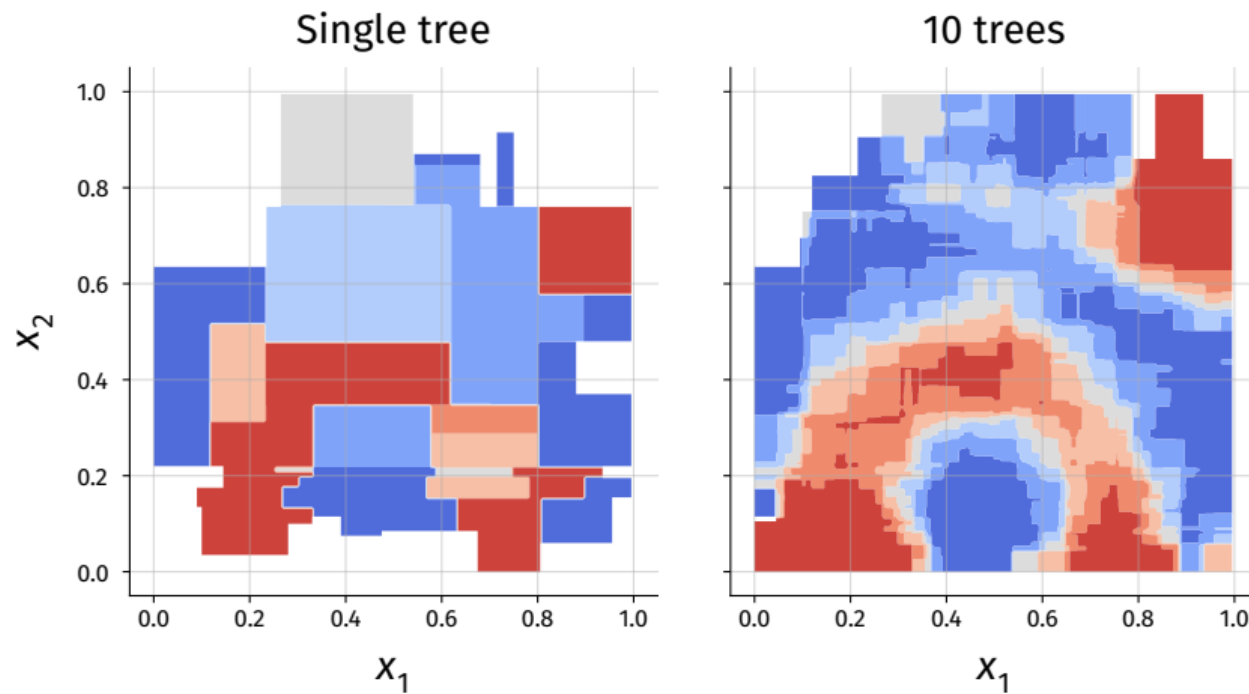
- Choose  $\delta \in [0, \frac{1}{2}]$
- Sample  $s$  in  $[a + \delta(b - a), b - \delta(b - a)]$
- $\delta = 0 \implies s \in [a, b]$  (uniform)
- $\delta = \frac{1}{2} \implies s = \frac{a+b}{2}$  (centered)



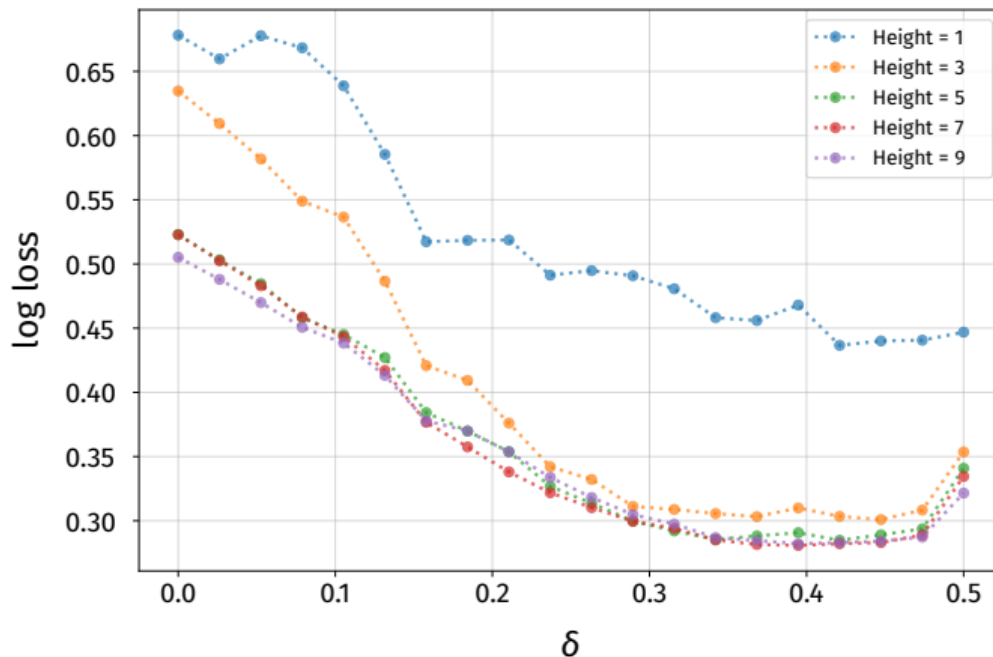
# Some examples



# Banana dataset with $\delta = 0.2$



# Impact of $\gamma$ on performance



# Tree regularisation

- A decision tree overfits when its leaves contain too few samples
- There are many popular ways to regularise trees:
  1. Set a lower limit on the number of samples in each leaf
  2. Limit the maximum depth
  3. Discard irrelevant nodes after training (pruning)
- None of these are designed to take into account the streaming aspect of online decision trees

# Hierarchical smoothing

- Intuition: a leaf doesn't contain enough samples... but its ancestors might!
- Let  $G(x_t)$  be the nodes that go from the root to the leaf for a sample  $x_t$
- **Curtailment** [ZE01]: use the first node in  $G(x_t)$  with at least  $k$  samples
- Aggregated Mondrian trees [MGS19] use context weighting trees

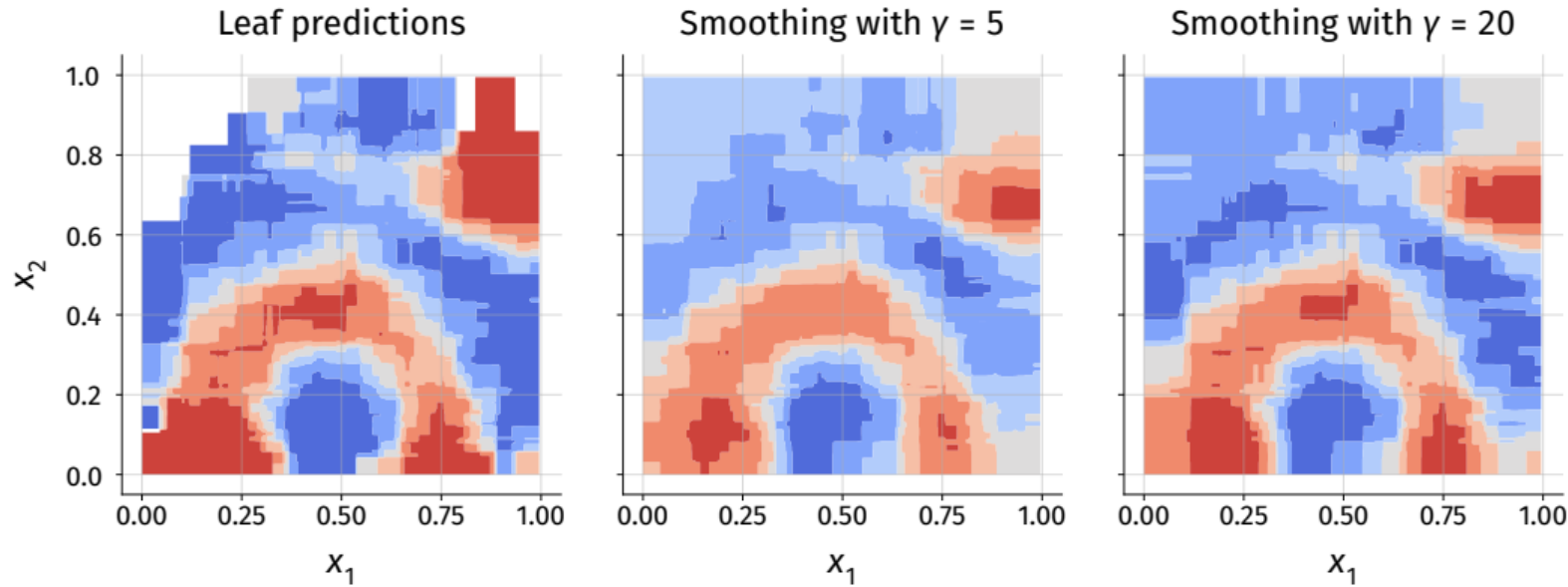
# A simple averaging scheme

- Idea: make each node in  $G(x_t)$  contribute to a weighted average
- Let
  - $k$  be the number of samples in a node
  - $d$  be the depth of a node
- Then, the contribution of each node is weighted by:

$$w = k \times (1 + \gamma)^d$$

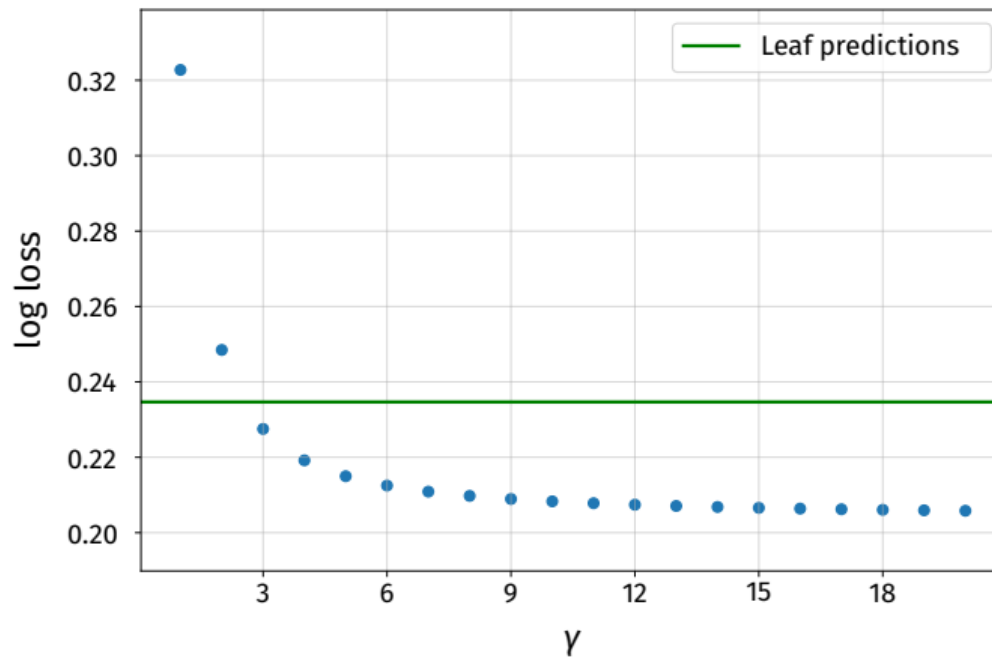
- The more samples a leaf contains, the more it matters
- The deeper a leaf is, the more it matters
- $\gamma \in \mathbb{R}$  controls the importance of both values
- I like to call this **path averaging**

# Averaging on the banana dataset



Notice what happens in the corners.

# Impact of $\gamma$ on predictive performance



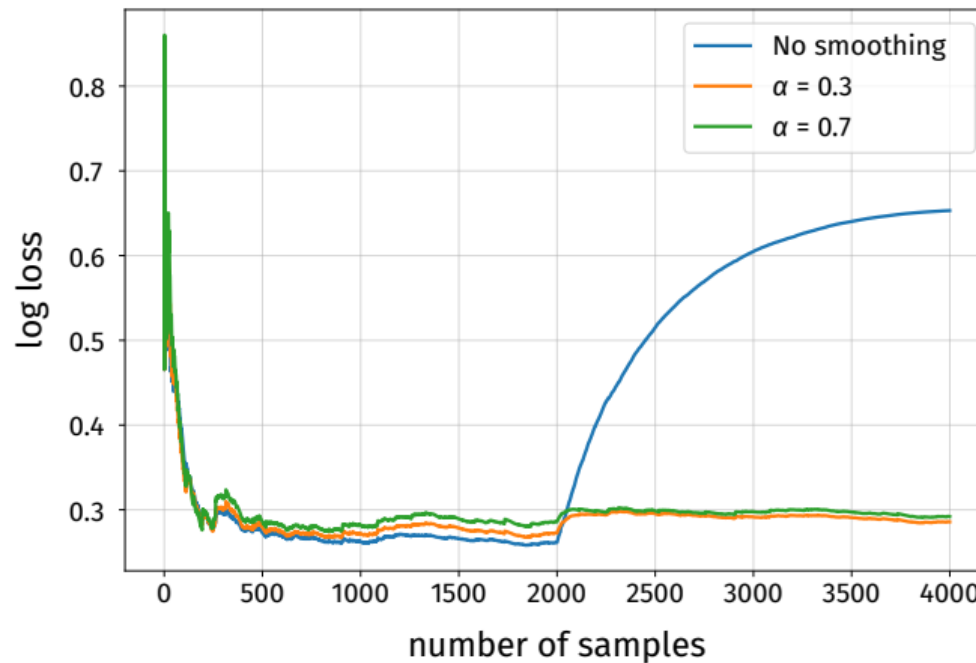
# Dealing with concept drift

- Each node contains a running average of the  $y$  values it has seen
- Instead, we can maintain an exponentially weighted moving average (EWMA):

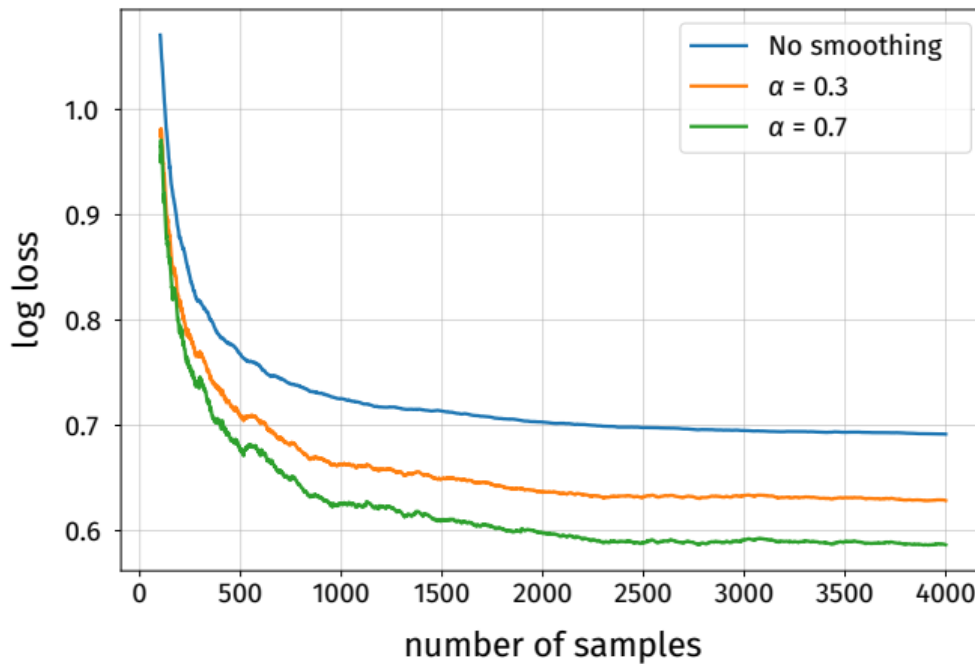
$$\bar{y}_t = \alpha y_t + (1 - \alpha) \bar{y}_{t-1}$$

- $\alpha$  determines the influence of the most recent values

# Hard drift: flip y values after 2000 samples



# Soft drift: slowly rotate samples around barycenter



# Feature selection

*Final paragraph from [MGS19] A limitation of AMF, however, is that it does not perform feature selection. It would be interesting to develop an online feature selection procedure that could indicate along which coordinates the splits should be sampled in Mondrian trees, and prove that such a procedure performs dimension reduction in some sense. This is a challenging question in the context of online learning which deserves future investigations.*

Online feature selection is a difficult problem!

# A solution?

1. Initially, we don't know the importance of each feature, so we pick them at random
2. After some time, we can measure the quality of each split within each tree
3. We can derive the feature importances from the splits each feature participates in
4. Every so often we can build a new tree by sampling feature relative to their importances
5. The selection probabilities should be conditioned on the features already chosen

This is still work in progress, but there is hope.

# Parameters recap

- $m$ : number of trees
- $h$ : height of each tree
- $\delta$ : the amount of padding
- $\gamma$ : determines how the path averaging works
- $\alpha$ : exponentially weighted moving average parameter

# Some useful Python libraries

- `scikit-garden` – Mondrian trees
- `onelearn` – Aggregated Mondrian trees
- `scikit-multiflow` – Hoeffding trees
- `scikit-learn` – General-purpose batch machine learning
- `creme` – General-purpose online machine learning

# Train/test benchmarks

	Moons	Noisy linear	Higgs	Higgs*
Batch log reg	.324	.244	.640	.677
Batch log reg with Fourier features	.193	.213	.698	.641
Batch random forest	.225	.210	.615	.639
NN with 2 layers	.171	.196	.653	.637
Online log reg	.334	.323	.662	.677
Mondrian forest	.349	.316	.692	.905
Aggregated Mondrian forest	.205	.199	.671	.649
Hoeffding forest	.330	.258	.664	.649
Padded trees (us)	.185	.193	.678	.644

# Streaming benchmarks

Work in progress!

*Slides are available at  
[maxhalford.github.io/slides/online-decision-trees.pdf](https://maxhalford.github.io/slides/online-decision-trees.pdf)*

*Feedback is more than welcome.*

*Stay safe!*

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